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PY, ex 2

(54) Drug delivery system

(57) Particles of drug are directed away from the reticuloendothelial system by use of a surface coating which prevents the take up of the composite particles by the liver. Suitable coating agents are Tetronic (RTM), poloxamers, polysaccharides, xanthan, hyaluronic acid etc.

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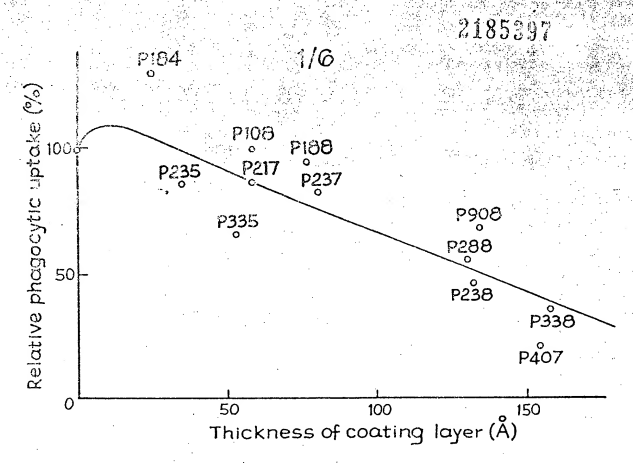
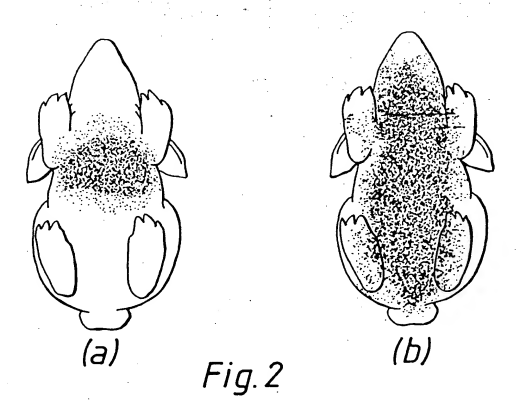


Fig.1



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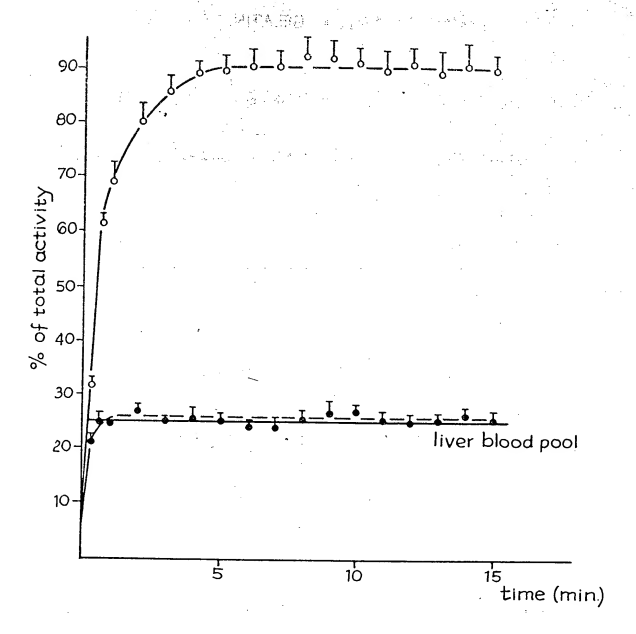
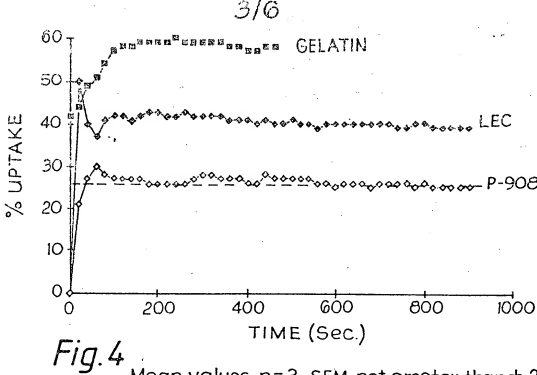
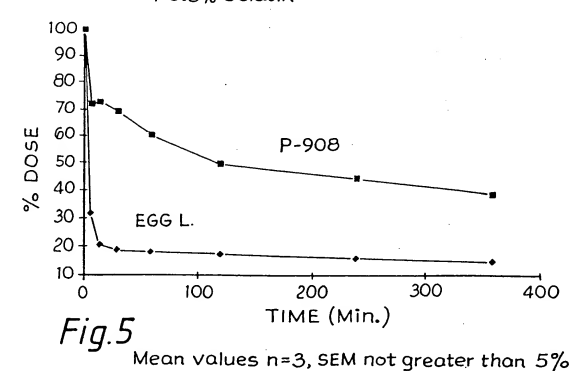


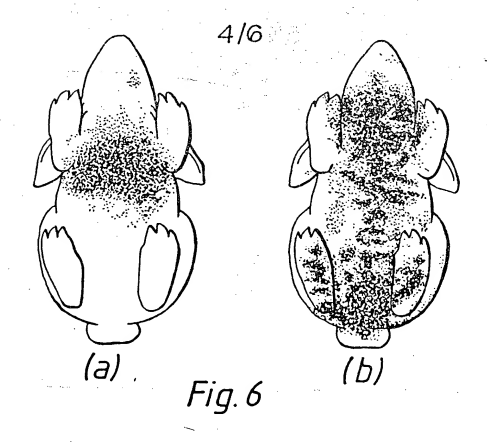
Fig. 3

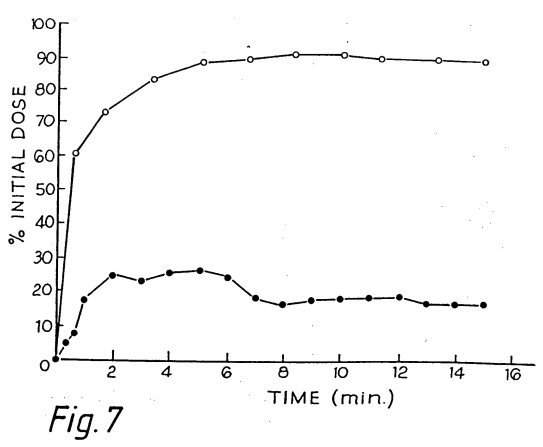


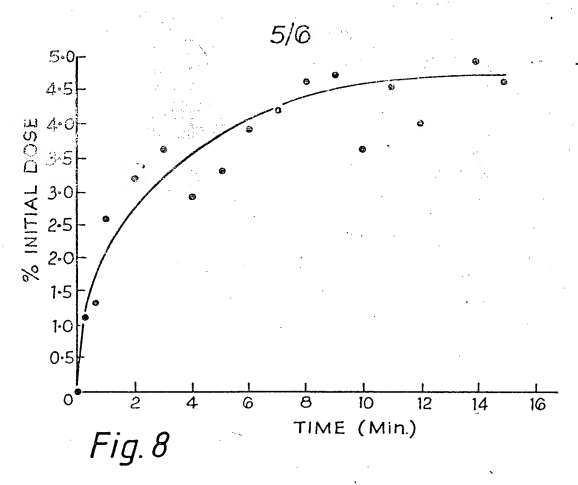
Mean values n=3, SEM not greater than $\pm 2\%$ Dotted line at 25% Indicates blood pool.

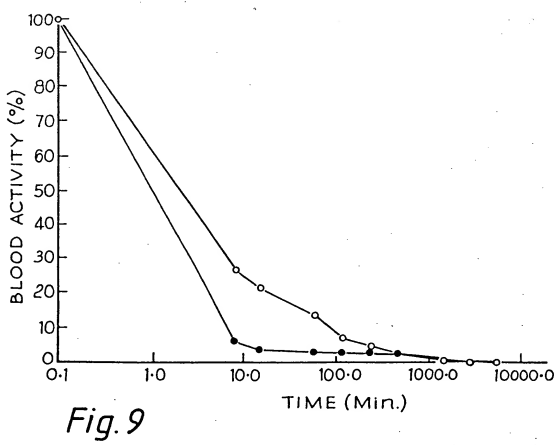
Values at 6 hours:	% Uptake in liver
1.2% Lecithin	34±2
1%	27 <u>+</u> 2
1.2% Lecithin	47 <u>+</u> 1



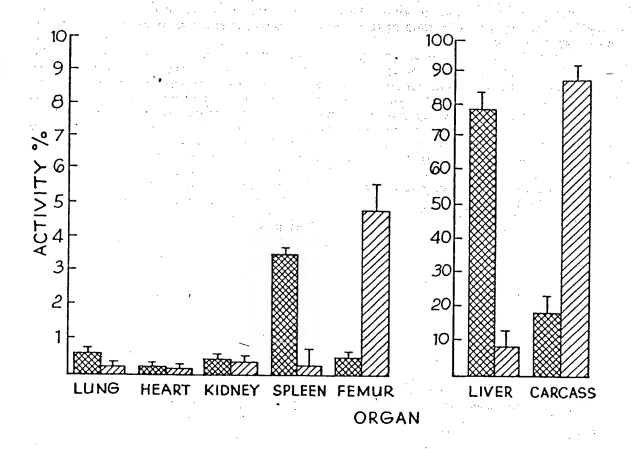








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KEY:

- $(oxine{oxtimes})$ Uncoated microspheres
- (🖾) Poloxamer 407 coated microspheres

Fig. 10

SPECIFICATION 1397 programme and officer and

or a many free case, edg. 1, 1, 1, 160 (1997). Drug delivery system

The present invention relates to drug delivery systems and more pain related to a system for assisting in the delivery of a drug or radiodeagnostic agent to a desired location within the animal or human body.

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been proposed as a means of directing drugs contained therein to specific sites in the body. 10. This concept, also known as drug targeting, has been well described in a number of publications, review articles and books. (see for example Davis, Illum, Tomlinson and McVie, (editors) Microspheres and Drug Therapy, Elsevier, Amsterdam, 1984). Colloidal carriers have been shown to perform well in vitro tests but their utility in vivo has been disappointing. It is known to be a relatively simple matter to direct particles to the lung or to the liver by exploitation of physical 15 factors such as particle size. However, the rapid and efficient capture of injected particles by the cells of the reticuloendothelial system residing in the liver (namely the Kupffer cells) does present

Colloidal particles in the form of microspheres, microcapsules, emulsions and liposomes, have

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a major obstacle to targeting colloidal particles elsewhere. Indeed, in a recent review article by Poste and Kirsch (Biotechnology 1: 869, 1984) and Posnansky and Juliano (Pharmacol. Revs. 36,277, 1984) this very point was emphasised. Similarly, at a meeting of the New York 20 Academy of Science held in March 1984 (published in Proceedings of the New York Academy

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of Sciences, Vol. 446, Editors Tirrell, D.A., Donaruma, L.G. and Turek, A.B., 1985), on the topic of polymers for drug delivery, many of the presenters of papers concluded that it would be almost impossible to direct colloidal particles to other sites than the liver and spleen when administration was by the intravaneous route. The present invention provides a method whereby 25 it is possible to direct particles away from the reticuloendothelial system residing in the liver and

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spleen by the use of surface coatings (and surface grafting techniques).

The State of the State of the

Model particles for use in studying the fate of drug carriers are often used in order to determine the scientific basis of drug targeting. Polystyrene microspheres of different sizes have been particularly useful in this respect. The small polystyrene particles of a size less than 100 30 nm are administered intraveneously. They are taken up rapidly and efficiently in the liver as measured by the non-invasive technique of gamma scintigraphy or by studies on animals where organs are removed and radioactivity levels are determined in such organs. Typically, more than 90% of the injected dose is found within the liver in a period of about 3 minutes (Illum, Davis, Wilson, Frier, Hardy and Thomas, Intern. J. Pharmaceutics 12 135 (1982)).

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It is an object of the present invention to provide a drug delivery system which obviates the above problem and prevents such a rapid take up of any injected dose by the liver.

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According to the present invention there is provided a drug delivery system comprising a number of particles containing an active drug, or a diagnostic agent to include radioactive materials. The particles could be for example, emulsions, mirospheres made from natural and synthetic polymers, or phospholipid vesicles, each particle being coated with a material to form a 40 composite particle which substantially prevents the take up of the composite particle by the liver.

Preferably the particles are coated with a material that provides them with both a hydrophilic coat that will minimize the uptake of blood components and a steric barrier to particle-cell 45 interaction. It is then found that the amount being taken up by the liver is greatly reduced. One preferred material is the block copolymer known as tetronic 908. This is a non-ionic surfactant which is obtained by polycondensation of propylene oxide and ethylene oxide on ethylenediamine. This coating material allows intravenously injected particles to remain within the systemic circulation with minimal uptake in the liver and spleen. Another preferred material is the block 50 copolymer known as poloxamer 407, a mixture of polyoxyethylene and polyoxypropylene domains. This material also is effective at preventing uptake of coated particles in the liver and spleen but directs them almost exclusively to the bone marrow. Other members of the poloxently large hydrophilic domain for steric stabilization. Typically an adsorbed layer thickness of 55 about 100 Angstrom or larger is required. This represents in the poloxmer series 60 or more

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amer and poloxamine series have similar effects provided that the material chosen has a sufficiethylene oxide units.

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The mechanism of action of the materials resides in the structure of the coating agent, namely that it has hydrophilic and hydrophobic domains. The hydrophobic domain will anchor the coating to the particle surface and prevent its displacement by plasma proteins. A suitable 60 molecular weight for this domain will be 4000-5000 Daltons. Hydrophobic domains include polyoxypropylene groups as well as other hydrophobic moeities that can be incorporated into polymer chains. For example, esterified maleic acid groups.

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The hydrophilic domain should be of a sufficient size and hydrophilic nature to prevent (or at least minimise) the coating of the particle by blood components (that is to minimise the pheno-65 menon known as onsonisation) as well as to provide a steric barrier so as to provide etoric

	stabilisation, a phenomenon well known in the field of colloid science (Napper, Polymeric Stabilisation of Colloidal Dispersions, Academic Press, London, 1983). Such steric stabilisation serves to prevent the interaction of particles with the macrophage cells of the reticuloendotheilial	
	5 Daltons. Embodiments of the present invention will now be described, by way of example with reference to the accompanying drawings in which:	· 5
1	polystyrene particles; so Carbota, and the second polystyrene particles and polystyrene particles. So Carbota, and the second polystyrene particles are carbotal to the second polystyrene particles.	
	and poloxamine 908-coated (b) polystyrene particles (60 nm); where the stration of uncoated (a) Figure 3 shows activity-time profiles for the uptake of uncoated (3) and coated (908) (O) particles in the liver (n=3, mean ± SEM); where the strategy of the	÷
1	Figure 6 shows gamma camera scintiscans of rabbits 3 hours after intravenous administration of 131,-labelled polystyrene microspheres (60 nm) (a) uncoated (b) poloxamer 407-coated; Figure 7 shows activity profiles for liver/spleen region after administration of 131,-labelled polystyrene microspheres (60 nm) (a) uncoated (b) poloxamer 407-coated;	15
20	O Figure 8 shows a graph of uptake of poloxamer 407 coated microspheres in the hind leg of the rabbit as measured by gamma scintigraphy:	20
_	Figure 9 shows a graph of the activity in the circulating blood after the administration of 131,—labelled polystyrene microspheres uncoated, O poloxamer 407 coated; and Figure 10 shows the distribution of 131, labelled polystyrene microspheres in various organs 8	
2!	microspheres.	25
30	Practical studies conducted in vitro with serum on the uptake of coated and uncoated particles by mouse peritoneal macrophages have demonstrated the importance of anchoring the polymer coating to the surface of the particle and surface layer thickness.	•
	Surface layer thickness Polystyrene particles (60 nm in diameter) were dialysed against distilled water for 3 days. 4.0% w/v aqueous solutions of the various polyamers and polyamers and polyamers.	30
35	measurements, remained above the plateua level of the adsorption isotherm i.e. above the critical micelle concentrations. Aliquots of 2.5% w/v polystyrene particles and the coating solution were mixed and incubated at room temperature overnight. The particle suspension was then diluted with distilled water (20 ul per 10.0 ml) and the pH adjusted with UCL or NOOL. The state of the coating solution were	35
40	particles at pH 2.1, 3.0, 5.5 and 9.5 using photon correlation spectroscopy.	40
45	Mouse peritoneal machrophage studies Polystyrene microspheres of 5.25 min diameter were chosen for the mouse peritoneal macrophage studies because uptake could be measured by a microscopic method and van der Waals attractive forces would be a dominant factor thereby allowing differentiation of the stabilising capacities of different block copolymers. The polystyrene microspheres were dialysed against distilled water for 3 days to remove any surfactant present. The particles were then incubated for 24 hours with the different 2% w/v poloxamer and poloxamine solutions. The	45
50	adsorbed material was in the plateau region of the respective adsorption isotherems. Female NMRI mice (Bommice, Monholtgaard Breeding and Research Centre Ltd., Ry, Denmark) weighing 20–25 g, were used to provide the peritoneal macrophages. The approach was tilled.	50
55	culture Medium El99 concentrate (10 x) (Flow Laboratories), 10 ml swine serum, 2.5 ml sodium bicarbonate 7.5%. 0.1 ml crystamycin, 6 mg heparin, 77.4 ml sterile water) injected into the peritoneal cavity followed by a smaller volume of sterile air. The peritoneal wall was gently massaged and the medium containing the macrophages was withdrawn and collected in a sterile	55
	and pooled. A cell count was conducted using a Coulter Counter (model TAII). The viability of the macrophages was tested by exclusion of tryptan blue and found to be in the order of 95%. The macrophage suspension was adjusted to a final cell count of 1.0 × 10 ⁶ cells/ml and 1.25 ml of this suspension pipetted into each 30 mm dish to give 1.25 × 10 ⁶ cells per plate. The plates were incubated at 37°C in 95% air/5% CO ₂ for 3 h to permit machrophage adherence to the	60
65	washed once with sterile PBS. 1.25 ml of cell culture medium added (10 ml Modium 5100).	^= \frac{1}{2}

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concentrate (10 X), 10 ml Medium E199 concentrate (10 X), 10 mlswine serum, 2.5 ml sodium bicarbonate, 0.1 ml crystamycin, 10 mg L-glutamine and 78.8 ml sterile water), and the plates incubated at 37°C in 95% air/5% CO₂ for 24 h. After incubation the medium was removed and of the cells washed once with sterile PBS. Then 2.5 ml cell culture medium containing the appropriate number of coated or uncoated micropheres (5 particles per macrophage) was added to each plate and the plates incubated in groups of 3 for 15, 30, 45, 60 and 90 min, as determined beforehand by a time course experiment. Before counting the number of particles phagocytosed by the macrophages, the media was removed from the plates, the cells washed 2 times with sterile PBS and fixed with methanol for 5 min. Then the cells were stained with Giernsa (1:10) for 15 min and washed with water. The plates were left to dry and the number of microspheres phagocytosed by the macrophages was counted for a total of 100 macrophages using a light microscope at a magnification of 500 times. The experiments were performed in triplicate and results were expressed as the number of microspheres phagocytosed by a 100 macrophages.

Experiments were also performed to determine whether free poloxamer and poloxamine had any effect on the ability of the macrophages to phagocytose particles. 2% w/w aqueous solutions of the coating agents were added to the cells and left to incubate 1 hour. The solution was removed and the cells washed 2 times with PBS. Then the cell culture medium containing the unocated microspheres was added and the degree of phagocytosis determined as before. Free polymer was found not to influence phagocytosis and therefore in all experiments the excess poloxamer of poloxamine was not removed before incubation with macrophages.

The relative uptake of the various coated 5.25 µm polystyrene particles by mouse peritoneal macrophages and the relationship with surface layer thickness are shown in Table 1 and Fig. 1. In general terms it can be seen that the greater the absorbed layer thickness the lower the relative phagocytic uptake. These results are in line with the predictions of the various theories put forward to explain the phenomenon of steric stabilisation. Therefore, it appears that these theories can also be applied to the interaction of particles with phagocytic cells. Extrapolation of the regression line shown in Fig. 1 to zero phagocytic uptake predicts that an adsorbed layer thickness of about 230 A would be necessary to overcome van der Waals attractive forces between macrophages and 5.25 µm particles.

The size of the layer that would be sufficient to give the same stabilising effect for much

The size of the layer that would be sufficient to give the same stabilising effect for much smaller (e.g. 60 nm) particles is difficult to predict exactly. However, since the van der Waals attractive forces (VA) are directly related to particle radius (a)

$$35 V_{A} = \frac{a A_{eff}}{12 h}$$

where A is the composite Hamaker constant and h is the Planck's constant, we would expect that an adsorbed layer thickness of about 100 A should be adequate to provide not only steric stabilisation of 60 nm polystyrene particles in terms of their aggregative propensity but also to a . 40 lack of interaction with macrophages.

Embodiments of the present invention will now be described, by way of examples:-

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EXAMPLE 1

45 THE ORGAN DISTRIBUTION AND CIRCULATION TIME OF INTRAVENOUSLY INJECTED COLLOI-DAL CARRIERS STERICALLY STABILIZED WITH A BLOCK COPOLYMER—POLOXAMINE 908 Methods

Polystyrene microspheres in the size range 50-60 nm were obtained from Polyscience (Northampton, UK). The particle size was confirmed using photon correlation spectroscopy. The particles were surface labelled with lodine-131 as described previously by L. ILLUM, S.S. DAVIS, C.G. WILSON, N.W. THOMAS, M. FRIER and J.G. HARDY, Int. J. Pharm. 12, 135, 1982).

Poloxamine 908 (average Mw 25000: 80% average weight percentage of polyoxyethylene chains) was obtained from Ugine Kuhlman Ltd., Bolton UK and used as received.

Incubation of the polystyrene microspheres with a 2% w/v solution of poloxamine 908 gave 55 an adsorbed layer thickness of 134 A.

In vivo experiments were conducted with groups of New Zealand White rabbits (3 kg) (n=3). Intravenous injections were given via the marginal ear vein (polystyrene microspheres 0.3 ml, 4×10^{13} particles, 3 MBq activity; emulsions 1.0 ml, 10^{12} particles, 3-4 MBq. Uncoated polystyrene particles were administered in distilled water (control). Particles coated with poloxamine 908 (24 hours equilibrium) were administered either as the incubation mixture (containing 1% poloxamine 908) or in distilled water after the excess poloxamine had been separated on a Sepharose CL4B column.

One group of rabbits was given similar repeated injections of poloxamine coated polystyrene microspheres on five consecutive days. Another group was given a dose of uncoated polystyrene microspheres 1 hour after the injection of the coated material and the coated material.

i			
	<u> </u>	Blood samples were taken at suitable intervals and the activity counted in a gamma counter. The distribution of the labelled particles in the liver were followed by gamma scintigraphy. Dynamic and static images of the liver distribution were analysed by creating regions of interest and compared to whole body activity. The activity in the liver associated with the blood pool was determined to be 25% of circulating activity using sequential administration of Tc-99m labelled pyrophosphate (red blood cell label) and lodine-131 labelled microspheres (to provide a	5 5
	10	and C. McMARTIN, J. Endocr. 82, 33, 1979), J.W. TRIPLETT, T.L. HAYDEN, L.K. McWHORTER, S.R. GAUTAM, E.E. KIM and D.W.A. BOURNE, J. Pharm. Sci. 74, 1007, 1985). Eight days after administration the rabbits were sacrificed and organs removed. Total activity in selected sites and in the carcass was determined using a large saple volume gamma counter.	10
	15	Uncoated polystyrene particles were taken up rapidly (t _{50%} =55 s) and efficiently (90% of dose in 2 min) by the liver and spleen while particles coated with poloxamine 908 remained largely in the vascular compartment and demonstrated little uptake in the liver/spleen region (Figs. (2-3). Similar results were obtained for coated particles separated from excess polyamine 200.	15
	20	sepharose CL4B column. Repeated injections of polystyrene particles coated with poloxamine 908 (one injection per day for 5 days) resulted in some uptake in the liver and spleen, but this was largely associated with the blood pool in the liver. The injection of uncoated polystyrene particles into rabbits 1 hour after they had received a does of polystyrene particles coated with poloxamine 908 demonstrated that the uncoated particles were mainly removed by the liver/spleen as for untreated animals whereby demonstrating that the poloxamine 908 had caused no	- 20
	. 0-	iniparment of the reticuloendothellal system.	
	25	largely in the vascular compartment while in correspondence with the scintigraphic information, little of the uncoated material could be found in the blood (Table 2). Interestingly, a significant	25
, other s	30	Scintigraphic measurements and organ level determinations (see below) failed to reveal significant sites of uptake (including bone marrow). Consequently, it is suggested that the coated particles could be loosely associated with endothelial cells lining the vaculature. Levels of activity in the different organs eight days after injection that are shown in Table 3. The uncoated particles were found largely in the liver and in the spleen while the coated	30
	.35	particles were largely associated with the carcass.	
		EXAMPLE 2	35
		INTRAVENOUS ADMINISTRATION OF RADIOLABELLED EMULSIONS AND THE ROLE OF THE BLOCK COPOLYMER—POLOXAMINE 908	
	40	This study was performed in order to establish whether the coating agent poloxamine 908 would retain a biodegradable emulsion system solely within the systemic circulation. Emulsions labelled with the gamma emitting agent iodine-123, were injected intravenously into rabbits. Two control formulations consisted of emulsions prepared using egg lecithin as the emulsifier with	40
	45	that gelatin can have an important role in directing colloidal particles to the liver; the process being mediated by the absorption of the blood component fibropactin. The role of the difference	45
	3	scintigraphic imaging of the livers of rabbits over a suitable period of time, as well as the removal of blood samples and the counting of gamma activity. The oil chosen for this week was to be considered by	-
	50	material is metabolised by the body, scintigraphic and blood level data were collected over a period of 6 hours.	.50
	55	Methods Animals Female New Zealand White rabbits of an approximate weight of 2 has a	55
		experimental model, 3 rabbits were chosen per group. Preparation of emulsions	- 55
i	b	Soybean oil was labelled using the method of Lubran and Pearson (J. Clein. Pathol. 11 (165) (1985). Iodine-123 was chosen as the most suitable radio-nuclide from the standpoint of its good imaging characteristics, its short half life and its greater safety over iodine-131. The iodine-123 was obtained from Harwell. The iodination method involves the covalent attachment of	60
đợ.		small quantities of labelled iodine across the double bond of the unsaturated components of the vegetable oil. This method has been used with success previously and similar iodinated fatty acids have been used in the radio-diagnostic field as myocardial imaging agents. The radio-	65

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labelled oil was mixed with a further proportion of unlabelled oil and the mixed oil was then emulsified with either poloxamine 908 (6ASF) (2%) or with egg lecithin (Lipoid) (1.2%). An ultrasonic probe system (10 min sonication) (Dawe Soniprobe) was employed for this procedure. Previous investigations using unlabelled oils has indicated that the particle size produced by this method was of the order of 150 mm. This size is very similar to that found in commercial fat emulsion products (e.g. Intralipid). One sample of the egg lecithin stabilised emulsion was mixed with gelatin (2%) according to the procedure described by Tonaki et al (Exp. Mol. Path. 25 189 化重点燃料 化铁油 化抗压性 化氯磺基 (1976)....

In this process some of the gelatin is adsorbed onto the surface of this particles or may form 10 a mixed emulsifying layer with the egg lecithin and will thereby potentiate uptake of the emulsion in the liver, mediated by adsorbed fibronectin.

Experimental procedure

The experimental animals were injected via the marginal ear vein using 1 ml samples of the 15 labelled emulsions. The oil content in the emulsions was 10%. The emulsions were followed by a 2 ml flush of normal saline. Following injection the animals were placed on the measuring surface of a gamma camera (Maxi camera, GEC, 40 cm field of view) tuned to the photoenergy peaks of iodine-123. Dynamic images were taken every 15 seconds over a period of 15 minutes. Blood samples were removed from the contralateral ear (0.5 ml). The scintigraphic 20 images were stored on computer and then analysed to provide information of the liver (spleen) uptake. Blood samples were diluted and counted in a conventional gamma counter. It is noted here that with gamma scintigraphy it is difficult to distinguish between the liver and spleen in a live animal but, with reference to Fig. 10 and to other results it is the liver which is the dominant organ. 25

Results

Uptake of labelled emulsions in the liver and spleen region is shown in Fig. 4. It can be seen that the extent of uptake is dependent upon the nature of the emulsifier used in preparing the emulsions. Those prepared using poloxamine 908 provided a liver uptake of approximately 25% 30 while those emulsified with egg lecithin had a value closer to 40%. The emulsions containing the added gelatin had an uptake value of approximately 60%. These liver uptake values for egg lecithin and P-908 systems are reflected in the blood level versus time profile in that the emulsions stabilised by egg lecithin demonstrate a much faster clearance from the blood than thoe stabilised by poloxamine 908 (Fig. 5). The rapid fall in blood level seen for both curves can 35 be attributed to the presence of the small quantities of free iodine that was administered. A kinetic analysis of the data (first order) indicates that the egg lecithin stabilized emulsion is cleared from the blood with a half life of about 5 mins while the P-908 stabilized emulsion is cleared with a half life of about 208 minutes. The plateau level of activity seen for the egg lecithin data reflect the fact that the emulsion is being metabolised and iodinated breakdown

40 products are being released into the plasma to give a more or less steady state level. The activity recorded in the liver of an animal after the administration of a colloidal system will include activity resulting from the uptake of those particles by liver cells (most probably the Kupffer cells) as well as normal circulating activity as part of the blood pool. This approximates to 25%. Thus in the studies conducted with poloxamine 908 it can be concluded that all the 45 activity recorded in the liver (spleen) region is due to circulating unsequestered emulsion and that the block copolymer effectively prevents liver uptake of the emulsion.

The results of the study confirm the investigations conducted by using polystyrene microsphere's coated with the block copolymer poloxamine 908 that such systems are largely ignored by the liver and are kept in circulation for an extended period of time. Such systems could have 50 great advantages for the delivery of pharmacological agents, where uptake of emulsion particles by the liver needs to be avoided to prevent adverse reactions and side effects.

EXAMPLE 3

TARGETING OF COLLOIDAL PARTICLES TO THE BONE MARROW USING THE BLOCK COPO-55 LYMER—POLOXAMER 407

The purpose of this study was to evaluate the extent and site of the diversion of the poloxamer 407 coated polystyrene particles in the intact animal model. This material has the ability to deliver model colloidal particles selectively to the bone marrow.

60 Methods

Polystyrene particles (60 nm in diameter) were purchased from Polyscience (Northampton, UK). The particle size was confirmed using photon correlation spectroscopy (PCS). The particles were surface labelled with iodine-131 was described previously. Poloxamer 407 (average MW 10500) was provided by Ugine Kuhlman Ltd., Bolton, UK, and used as received.

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poloxamer 407 providing a surface coating layer of 123 Å thickness as measured by PCS. Groups of New Zealand White rabbits (3 kg) (n=3) were injected intravenously via the marginal ear vein with either uncoated polystyrene particles (0.3 ml, 4×10^{13} particles, 3 MBg activity of particles coated with poloxamer 407 (0.6 ml, 4×10^{13} particles, 3 MBq activity). 5 Particles coated with poloxamer 407 were administered as the incubation mixture, uncoated particles in distilled water. The factors of war, notes to appropriate transplanting 化基础的概念 机红色 数数 Blood samples were taken at suitable intervals and the activity measured using a gamma counter. The distribution of the labelled particles in the body was followed by gamma scintigraphy. Dynamic and static images of the liver, spleen region and the left hind leg were analysed 10 by creating regions of interests and compared to the whole body activity. Eight days after 10 administration the rabbits were sacrificed and organs removed. Total activity in selected organs, blood, femur and remaining carcass was determined using a large sample volume gamma counter. 15 Results 15 Gamma camera scintiscans of the rabbits clearly demonstrated that uncoated polystyrene particles were largely taken up by the liver and spleen after injection while the poloxamer 407 coated particles were deposited in the bone marrow thereby providing a distinct picture of the rabbit skeleton. Furthermore, no images of the liver/spleen region or other organ regions could 20 be visualized. (Fig. 6). 20 The uptake of the uncoated particles by the liver/spleen region occurred both rapidly and efficiently with 90% of the particles being deposited in these organs within 2 min. This is illustrated in the liver/spleen activity-time profiles for the first 15 min after injection (Fig. 7). The poloxamer 407 coated particles showed a markedly decreased liver/spleen activity that reached a maximum of 25% after 2 min and then gradually decreased to a level of 17%. About 10% of 25 this activity can be attributed to the activity in the circulation (blood pool) and does not represent particle removal. During the same time period the poloxamer 407 coated particles were rapidly accumulated in the bone marrow with a half life of uptake of about 2 min as seen in the activity-time profile obtained by creating a region of interest around the left hind leg (Fig. 8). In comparison only background levels of activity were recorded for the same region of 30 interest in rabbits receiving the uncoated particles. Measured blood level activities showed that both the uncoated and coated particles were rapidly removed from the blood-stream. The estimated half lives of blood clearance correspond quite well with the measured half lives of uptake in the liver/spleen and the bone marrow, respectively (Fig. 9). Organ levels measured eight days after administration of the particles show conclusively that 35 coating the particles with poloxamer 407 leads to a reduction in lung, spleen and liver uptake. But more importantly a dramatic increase in the bone uptake is indicated by measured activity in the femur and the remaining carcass (Fig. 10). 40 DRUG DELIVERY APPLICATIONS 40 The particles coated with polocamine 908 that are retained in the blood stream could be used to target to other sites in the micro-vasculature, for example to subsets in the bone marrow, the liver itself, heart, kidney, lungs and even to tumour cells if the tumour had a vasculature that allowed extravasation. This type of targeting is terms active targeting and requires the attach-45 ment of a suitable ligand to the particle or to its polymer coat. Suitable ligands include monoclonal antibodies or their fragments, apolipoproteins, sugars and lectins. Drugs that could be administered using parrticles coated with poloxamine 908 include antiinfectives (for example amphotericin), macrophage activating agents, antithrombotics, cardiovascular agents (for example prostaglandins) and anti-leukamia drugs. The particle coated with poloxamer 407 could be used to direct drugs and radiodiagnostic 50 50 agents to the bone marrow. These include immunosupressants (cyclosporin), peptide drugs such as colony stimilating factors and radio-isotopes for diagnostic purposes (e.g. iodine isotopes, While the example given refers mainly to a model non-degradable particle, polystyrene, the same concept should work equally well with particles that will biodegrade in the body. Examples 55 include albumin, gelatin, polyalklcyanoacrylates, polylactides, polyglycolides, polyhydroxybutyrates and their mixtures in the form of copolymers, it also includes emulsions and phospholipid vesicles. The coating agent does not necessarily have to be a block copolymer comprising polyoxyethylene-polyoxy-propylene groups as shown in the example. Other materials that would provide the same type of effect could be used. Examples include poloxamers, polymaleic acid, polymers that

are esterified to produce suitable hydrophilic and hydrophobic domains as well as natural materials such as polysaccharides and hyaluronic acid. Polymer coatings that provide not only a steric barrier but also an electrostatic barrier are also effective in diverting particles away from

65 the reticuloendothelial system and materials such as vanthan gum which ?

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hydrophilic chain but also charged carboxyl groups are a suitable starting point provided it could be attached well to the surface of the colloidal particle in question. Colloidal particles in the form of liposomes and emulsions could also be coated with similar types of material. The results also indicate that the polymeric material tetronic 908 and macromolecules with similar hydrophilic/hydrophobic dornains could also be used as soluble macromolecular carriers for drug molecules by direct linkage or through degradable spaces and linkages.

Attachment of suitable hydrophilic groups to particles have been achieved by surface grafting techniques either during the polymerisation process whereby the particle is produced initially, or by subsequent grafting methods involving energetic sources such as ultraviolet light and gamma 10 irradiation.

Polexamine 908 and Polexamine 407 (CFTA names) are also available commercially under brand names TETRONIC and PLURONIC (Registered Trade Marks) from the BASF WYANDOTTE Corporation 100 Cherry Hill Road, P.O. Box 181 Parsippany N.J. 07054.

15 TABLE 1

THE PROPERTY OF THE PROPERTY O

Surface characteristics and phagocytic uptake of polystyrene particles coated with non-ionic surfactants

20	• •		olecular verage v		Thickness of coating layer	Relative phagocytic uptake		20
	Coating agent	18 25	(in mole	•		·		
25		≠≠≠ EO	PO	EO	A	<u>%</u>		٥٦
	None		<u>-</u> .	_	.0	100.0		25
	Poloxamer 108	165 46	16	46	58	100.4		
	Poloxamer 184	5 to 13	30	13	24	129.3		
	Poloxamer 188		30	75	76	95.4	·	
30	Poloxamer 217		35	52	58	87.6		30
	Poloxamer 235	27	. 39	- 27	35	86.5		0.5
	Poloxamer 237		39	97	132	47.0		
	Poloxamer 288	122	47	122	130	56.5		*
	Poloxamer 335	38	54	38	53	66.7	•	•
· 35	Poloxamer 338	128	54	128	158	36.7		35
	Poloxamer 407	98	67	98	154	21.6		
	Poloxamer 908			· <u>-</u>	134	69.5		

TABLE 2

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Blood Level Activity 15 mins and 1 hour after Administration of Uncoated and Coated Polystyrene Microspheres to Rabbits

	percentage of initial dose 15 min	e in blood (±SEM)
Polystyrne microspheres (PM) PM coated with	4.0 (±0.4)	3.0 (±0.01)
poloxamine 908	65.5 (±4.1)	60.0 (±4.1)
	microspheres (PM) PM coated with	Polystyrne microspheres (PM) 4.0 (±0.4) PM coated with

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	TABLE 3						
5	Deposition of Uncoated and Coated Polystyrene Microspheres in the Various Organs 8 days after Intravenous Administration in Rabbits. The Values are Expressed as Percentage of total Activity (±SE1)	5					
	Lung Heart Kidney Spleen Liver Carcass						
10	Polystyrene microspheres 0.15±0.01 0.11±0.01 0.22±0.02 1.45±0.20 59.5±6.9 38.6±7.1	10					
·. ·	(PN) A series of the first term of the series of the serie						
15	Poloxamer 338 0.51±0.03 0.22±0.01 0.34±0.03 0.93±0.19 30.2±5.5 67.9±5.7	15					
	PM coated with Poloxamine 980 2.50±0.80 0.20±0.01 1.50±0.20 1.20±0.10 18.9±3.2 73.7±2.4	•					
20		20					
25	CLAIMS 1. A drug delivery system comprising a number of particles of an active drug, each particle being coated with a material to form a composite particle which substantially prevents the take up of the composite particle by the liver.	25					
	 A drug delivery system as claimed in Claim 1 in which the particles are coated with a material that provides them with a hydrophilic coat and a steric barrier to particle-cell interaction. A drug delivery system as claimed in Claim 2 in which the coating material is the block copolymer known as tetronic 908. 						
30	4. A drug delivery system as claimed in Claim 1 in which the coating material is a poloxamer a polymaleic acid or a polymer that is esterified to produce suitable hydrophilic and hydrophobic	30					
35	5. A drug delivery system as claimed in Claim 1 in which the coating material is a natural material selected from the polysaccharides or hydronic acid.						
40	8. A drug delivery system substantially as described with reference to the accompanying	40					

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